



# Elemental stoichiometry and Rock-Eval® thermal stability of organic matter in French topsoils

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**Abstract.** The quality and quantity of soil organic matter (SOM) are key elements of soil health and climate regulation by soils. The Rock-Eval® thermal analysis technique is increasingly used as it represents a powerful method for SOM  
15 characterization by providing insights on bulk SOM chemistry and thermal stability. In this study, we applied this technique on a large soil sample set from the first campaign (2000–2009) of the French monitoring network of soil quality: RMQS. Based on our analyses on ca. 2000 composite surface (0–30 cm) samples taken all over mainland France, we observed a significant impact of land cover on both SOM thermal stability and elemental stoichiometry. Cropland soils had a lower mean value of hydrogen index (a proxy for SOM H/C ratio) and a higher thermal stability than grasslands and forests.  
20 Regarding the oxygen index (a proxy for SOM O/C ratio), we observed significant differences in values for croplands, grasslands and forests. Positive correlations between the temperature parameters on the one hand and the clay content and pH on the other hand highlight the protective effect of clay on organic matter and the impact of pH on microorganisms mineralization activity. Surprisingly, we found weak effects of climatic parameters on the thermal stability and stoichiometry of SOM. Our data suggest that topsoil SOM is on average more oxidized and biogeochemically stable in croplands. More  
25 generally, the high number and even repartition of data on the whole French territory allow to build a national interpretative referential for these indicators in surface soils.

## 1 Introduction

The fate of soil organic carbon (SOC) is crucial from both soil health and climatic perspectives. In terms of soil health, SOC plays an important functional role. Its decomposition by microorganisms provides energy to the whole soil food web and key  
30 nutrients to plants and soil fauna. SOC also regulates the water cycle through controlling soil structure (Rawls et al., 2003). From a climatic perspective, soils can act both as a source or a sink of carbon (Amundson, 2001; Eglin et al., 2010).



Maintaining or increasing SOC stocks has become a key policy issue for the coming decades (Rumpel et al., 2018) that raises a number of important scientific challenges regarding our knowledge of SOC dynamics (Dignac et al., 2017).

The evolution of SOC stocks depends on the balance between soil carbon inputs (mostly by plants) and outputs (mostly by microbial decomposition). The persistence of SOC determines soil carbon outputs so that estimating the biogeochemical stability of SOC to microbial decomposition (i.e. the difficulty for microorganisms to mineralize SOC) is of paramount importance to infer SOC dynamics (Schmidt et al., 2011; Lehmann and Kleber, 2015). Indeed, a better knowledge of SOC persistence would allow refining our estimates of the soil carbon inputs needed to maintain or enhance SOC stocks. However, estimating the biogeochemical stability of SOC is a challenging task because its turnover time encompasses a broad spectrum (ranging from some days or weeks to centuries; Balesdent and Guillet, 1982) resulting from a series of interacting SOC stabilization mechanisms. Indeed, SOC can be protected from microbial decomposition due to its chemical nature (e.g., pyrogenic SOC), its interactions with soil mineral surfaces or its spatial inaccessibility for microbes (Baldock and Skjemstad, 2000; Von Lützow et al., 2006).

Several routine techniques have been proposed to separate fractions that are labile, intermediate or stable at various timescales (von Lützow et al., 2007; Bispo et al., 2017; Chenu et al., 2015). However, none of these techniques manages to isolate precisely homogeneous fractions with the same biogeochemical stability (von Lützow et al., 2007; Poeplau et al., 2018; Cécillon et al., 2021). Common methods include biological respiration measurements performed during laboratory incubations of soils (e.g. Collins et al., 2000) and various physical (particle size or density) and/or chemical (aqueous or organic extraction) SOC fractionation methods (von Lützow et al., 2007).

Thermal analysis methods have been used for several decades to study the characteristics of soil organic matter (SOM). Many different methods exist, which measure different variables: e.g. thermogravimetry, differential scanning calorimetry, evolved gas analysis, etc. (Plante et al., 2009). A multitude of variations in temperature ramps, compositions of reaction atmosphere and measured parameters are encountered within each class of methods. Some thermal analysis methods provide indicators that are related to SOM biogeochemical stability: the more biogeochemically stable SOM is, the more thermally stable and energy- and hydrogen-depleted it is (Barré et al., 2016; Sanderman and Grandy, 2020). Among thermal analysis methods, Rock-Eval® thermal analysis is increasingly used to derive thermal indicators related to SOC biogeochemical stability (Gregorich et al., 2015; Saenger et al., 2015; Cécillon et al., 2018; 2021; Poeplau et al., 2019; Chassé et al., 2021).

The Rock-Eval® method was developed in the 1970s. Initially intended for the characterization of petroleum source rocks and sediments in order to estimate their potential for hydrocarbon extraction (Espitalié et al., 1977), this method was then adapted to the study of SOM (Disnar et al., 2003). This technique allows the measurement of the organic and inorganic carbon content of a soil sample, as well as numerous indicators of the thermal stability and elemental stoichiometry of SOM. Espitalié et al. (1977) has shown that the Rock-Eval® hydrogen index HI (respectively the oxygen index OI<sub>re6</sub>) is a good proxy for the H/C ratio (respectively O/C ratio) of organic matter in kerogens and, later, soils. Many temperature parameters can also be calculated to give insights on the thermal stability of SOM (Gregorich et al., 2015; Sebag et al., 2016; Cécillon et al., 2018). With a rate of one sample per hour at a reasonable price (below 50 USD per sample), Rock-Eval® thermal



analysis is a particularly fast and simple tool to use, and is therefore well suited to the study of large soil sample sets, such as the ones collected in the context of national or continental soil monitoring networks.

However, so far, the different existing soil monitoring networks worldwide have not used thermal analysis methods to infer SOC biogeochemical stability. Some of them have focused on SOC physical fractionation schemes, in combination with  
70 infrared spectroscopy or environmental variables (e.g. Vos et al., 2018; Viscarra-Rossel et al., 2019; Lugato et al., 2021; Sanderman et al., 2021). Here, we used Rock-Eval® thermal analysis to investigate the thermal stability and elemental stoichiometry of topsoil samples of the first campaign of the French national soil monitoring network (RMQS, Réseau de Mesures de la Qualité des Sols; GIS Sol; <https://www.gissol.fr/le-gis/programmes/rmqs-34>). The RMQS network has been  
75 designed for the long-term monitoring of the soil quality of the whole French territory by collecting information and sampling soils on, every 15 years in average, a set of 2170 sites at the locations of a regular, square grid thus forming a systematic sample (Jolivet et al., 2006; English version to be available online). The first campaign took place between 2000 and 2009 in mainland France, covering 7 major land cover types (croplands, grasslands, forests, vineyards & orchards, wastelands, poorly human-disturbed environments, and gardens).

In this study, we aimed in the first place to verify that the Rock-Eval® method was suited to characterize SOM on archived  
80 soil samples at the scale of a monitoring network. For this purpose, we checked if the organic and inorganic carbon yields of the Rock-Eval® thermal analysis for soil samples, calculated by comparing Rock-Eval® estimates to reference methods, were acceptable. Second, we computed several common Rock-Eval®-based indicators in order to perform an unprecedented country-wide evaluation of the thermal stability and elemental composition of the SOM. Third, thanks to the numerous environmental data available at each RMQS site, we aimed at studying the relationships between land-cover, climate and soil  
85 properties and the SOC-related indicators derived from Rock-Eval® thermal analysis.

## 2 Material and methods

### 2.1 Topsoil sampling and processing

A full description of the RMQS and of the soil sampling process of its first sampling campaign is available in Jolivet et al., 2006. Briefly, the soil is monitored at the locations of a regular, square grid with a resolution of 16 km. A sampling site was  
90 settled when possible at the center of the cell; otherwise, an alternative site was taken within a 1 km radius from the center of the cell. This resulted in a total of 2170 RMQS sites in mainland France. At each selected site, 25 topsoil samples (0–30 cm or tilled layer depths) were taken with a spiral soil auger from a 20 m × 20 m sampling area then mixed to provide a composite sample. Subsoil samples were also taken, but were not considered in the present study.

The composite samples (5 to 10 kg of bulk soil) were air-dried at 30°C in trays for 8 to 10 days on average. The samples  
95 were then quartered according to NF ISO 11464 to obtain a sub-sample of ca. 650 g. They were then crushed by hand to break aggregates while preserving calcareous and/or ferro-manganic nodules and sieved at 2 mm. The remains of the



composite samples were stored in water-tight plastic buckets. An aliquot of each air-dried and sieved composite sample was then finely ground using a Cyclotec 1093 (Foss).

Of the 2170 archived aliquots of finely ground topsoil samples from the first RMQS sampling campaign in mainland France, 100 2037 were recovered and used for this study. When necessary, the samples were manually ground again using an agate mortar to reach the particle size requirements of Rock-Eval® thermal analysis of soils (below ca. 250 µm).

## 2.2 Physical and chemical soil analyses

The physical and chemical soil analyses were carried out on the 2 mm sieved composite samples at the Laboratoire d'Analyse des Sols (INRAE, Arras, France). Among the large set of soil properties measured, we selected in this study the 105 following ones (Jolivet et al., 2006): particle-size measurements without decarbonation in g.kg<sup>-1</sup> of sample (Robinson pipette and underwater sieving, method validated in relation to standard NF X31-107) leading to five fractions (clay: ≤ 2 µm; fine silt: 2–20 µm; coarse silt: 20–50 µm; fine sand: 50–200 µm; coarse sand: 200–2000 µm); pH in water (NF ISO 10390, dilution with 1/5); total carbonate content in g.kg<sup>-1</sup> of sample (volumetric method NF EN ISO 10693) to estimate the total inorganic carbon  $C_{inorg} = \text{Total carbonate} \times 0.12$  in g.kg<sup>-1</sup> of sample; total carbon content (g.kg<sup>-1</sup> of sample) determined by 110 elemental analysis using dry combustion on non-decarbonated soil; organic carbon content derived from the elemental analysis (TOC<sub>ea</sub>; g.kg<sup>-1</sup> of sample) calculated as Total carbon -  $C_{inorg}$  (NF ISO 10694); total nitrogen in g.kg<sup>-1</sup> of sample (dry combustion NF ISO 13878); CEC in cmol+.kg<sup>-1</sup> of sample (cobaltihexammine chloride extraction NF X31-130); free iron oxides in g/100 g measured with the Tamm method in the dark and Mehra-Jackson method (INRA standard/NF ISO 22036).

## 115 2.3 Rock-Eval® thermal analysis

### 2.3.1 Thermal analysis process

Rock-Eval® thermal analyses were carried out at the UMR 7193 ISTeP (Sorbonne Université, Paris, France) on the 2037 recovered samples according to the routine classically used for soil samples (Disnar et al., 2003; Baudin et al., 2015). Approximately 60 mg of each finely ground topsoil sample was used for the Rock-Eval® thermal analysis on a RE6 turbo 120 device (Vinci Technologies, France). For each analysis, the sample was placed in a special high-temperature resistant stainless steel pod allowing the transport gas to pass through. It first underwent a pyrolysis step under an inert N<sub>2</sub> atmosphere. After a three-minute isotherm at 200 °C, the sample was heated to 650 °C following a temperature ramp of 30 °C/min. The flame ionization detector (FID) monitored the gaseous emissions of carbon from hydrocarbon compounds (HC\_PYR Rock-Eval® thermogram), while CO (CO\_PYR Rock-Eval® thermogram) and CO<sub>2</sub> (CO<sub>2</sub>\_PYR Rock-Eval® 125 thermogram) were detected by an infrared detector. The second step is an oxidation (laboratory air atmosphere with CO<sub>2</sub> and H<sub>2</sub>O previously removed, i.e., in presence of oxygen): the sample experienced a one-minute isotherm at 300 °C, then was raised to 850 °C following a 20 °C/min ramp, and finally remained on a five-minutes isotherm at 850 °C. The evolution of



CO and CO<sub>2</sub> were again monitored using the infrared detector during the oxidation phase (CO\_OX and CO<sub>2</sub>\_OX Rock-Eval® thermograms). The resulting five thermograms were processed using the Geoworks software (Vinci Technologies, 130 Geoworks V1.6R2), except for the parameters R-index, I-index which were computed using homemade Python scripts according to the formula proposed by Sebag et al. (2016).

Our Rock-Eval® thermal analyses campaign included duplicated soil analyses (one every eight samples), performed in order to check the reproducibility of the analyses, along with standard analyses (one every nine samples) to check the calibration of the device and identify a possible drift in the analysis. The Rock-Eval® thermal analysis of a soil sample measures its 135 total organic carbon (TOCre6) and total inorganic C (MinC) contents that sum to total carbon content (see Behar et al., 2001 for a detailed description). The organic carbon yield of Rock-Eval® thermal analysis was defined as TOCre6/TOCea, while its inorganic carbon yield was defined as MinC/Cinorg, and its total carbon yield was defined as (TOCre6+MinC)/(TOCea+Cinorg). We used the organic carbon yield of Rock-Eval® thermal analysis to select soil samples among duplicates: only the one with the best yield was conserved. When assessing SOM thermal stability and elemental 140 stoichiometry, it is essential to ensure that SOM analyzed by the thermal analysis method corresponds to SOM measured using the reference elemental analysis method. We therefore proposed a quality criterion for Rock-Eval® thermal analysis based on its organic carbon yield, with an arbitrary acceptable range of yields from 0.7 to 1.3.

### 2.3.2 Rock-Eval® parameters

Many usual Rock-Eval® parameters were calculated from the thermograms (Table A). First there are parameters related to 145 carbon quantities : the total organic carbon (TOCre6; g.kg<sup>-1</sup> sample); the total inorganic carbon (MinC; g.kg<sup>-1</sup> sample); the amount of pyrolyzable organic carbon (PC; g.kg<sup>-1</sup> sample); the ratio of pyrolyzable organic carbon over total organic carbon (PC/TOCre6; no unit); the carbon released during the first pyrolysis isotherm (PseudoS1; g.kg<sup>-1</sup> sample); the carbon released as hydrocarbons during the pyrolysis except during the first isotherm (S2; g.kg<sup>-1</sup> sample); the ratio of carbon released as hydrocarbons during the pyrolysis except during the first isotherm over the pyrolyzable organic carbon (S2/PC; no unit). 150 Second, there are temperature parameters related to the SOC thermal stability. Their calculation was performed over different intervals of integration depending on the thermogram. The upper limits of the integration ranges were selected to exclude CO and CO<sub>2</sub> signals derived from carbonates. The temperature parameter T50\_HC\_PYR (respectively T70\_HC\_PYR and T90\_HC\_PYR; °C) is defined as the temperature at which 50 % (respectively 70 % and 90 %) of the hydrocarbon effluents have been emitted during the pyrolysis ramp (the initial isotherm is excluded; the integration ends at 155 650 °C). Similarly, T30\_CO<sub>2</sub>\_PYR (respectively T50\_CO<sub>2</sub>\_PYR, T70\_CO<sub>2</sub>\_PYR and T90\_CO<sub>2</sub>\_PYR; °C) is the temperature at which 30 % (respectively 50 %, 70 % and 90 %) of the CO<sub>2</sub> have been emitted during the pyrolysis ramp (the beginning isotherm is excluded; the integration ends at 560 °C); T50\_CO\_PYR (°C) is the temperature at which 50 % of the CO have been emitted during the pyrolysis ramp (the beginning isotherm is excluded; the integration ends at 560 °C). T50\_CO<sub>2</sub>\_OX (respectively T70\_CO<sub>2</sub>\_OX and T90\_CO<sub>2</sub>\_OX; °C) is the temperature at which 50 % (respectively 70 % 160 and 90 %) of the CO<sub>2</sub> have been emitted during the oxidation phase (the integration ends at 611 °C); T50\_CO\_OX



(respectively T70\_CO\_OX; °C) is the temperature at which 50 % (respectively 70 %) of the CO have been emitted during the oxidation phase (the integration ends at 850 °C). We also calculated two other parameters previously used in assessing the thermal stability of SOC: the I-index (related to the thermolabile organic carbon released as hydrocarbon effluents, Sebag et al., 2016; no unit) and the R-index (the proportion of thermostable organic carbon released as hydrocarbon effluents after 165 400 °C, Sebag et al., 2016; no unit). Finally, we calculated the three following Rock-Eval® parameters, related to the SOM stoichiometry. The Hydrogen Index, HI, is the ratio of emitted hydrocarbons to TOCre6 (unit g HC.kg<sup>-1</sup> TOCre6); it is calculated following Eq. (1):

$$HI = \frac{S2 \times 100}{TOCre6} \quad (1)$$

where S2 is the hydrocarbons signal during pyrolysis (Behar et al., 2001). The Oxygen Index, OIre6, is the ratio of organic 170 oxygen to TOCre6 (unit g O<sub>2</sub>.kg<sup>-1</sup> TOCre6); it is calculated following Eq. (2):

$$OIre6 = \frac{16}{28} \times \frac{S3CO \times 100}{TOCre6} + \frac{32}{44} \times \frac{S3 \times 100}{TOCre6} \quad (2)$$

where S3 and S3CO are respectively the organic CO<sub>2</sub> and organic CO signals during pyrolysis (Behar et al., 2001; Cécillon et al., 2018). The ratio of hydrogen amount to oxygen amount is HI/OIre6 (no unit).

As presented above, the treatment of the five thermograms can result in the production of a multitude of Rock-Eval® 175 parameters. We have decided to present the results on the following parameters in more detail: T50\_HC\_PYR; T90\_HC\_PYR; T50\_CO2\_PYR; T50\_CO2\_OX; I-index; R-index; HI and OIre6. The results obtained for some other Rock-Eval® parameters are presented as Supplementary Information. The temperature parameters T90\_HC\_PYR, T50\_CO2\_PYR and T50\_CO2\_OX were selected as they are derived from the 3 different thermograms contributing the most to the Rock-Eval® signals, and as they are well correlated to the proportion of centennially stable SOC in temperate soils (Cécillon et al., 180 2021) and were used in some previous studies (e.g. Barré et al., 2016; Poeplau et al., 2019). The parameters T50\_HC\_PYR, I-index and R-index were selected as they were used in several previous studies (e.g. Gregorich et al., 2015; Sebag et al., 2016; Matteodo et al., 2018; Soucémariadin et al., 2018). HI and OIre6 were selected as they are usual Rock-Eval® parameters and they give insights on the elemental stoichiometry of SOM.

## 2.4 Climate data

185 Climate data were extracted from the French SAFRAN database (<https://publitheque.meteo.fr/okapi/accueil/okapiWebPubli/index.jsp>). The daily data were averaged over the 1969-1999 period in order to compute for each site the mean annual temperature (MAT) and mean annual precipitation (MAP).

## 2.5 Statistical analysis

We calculated linear regressions without intercept using the measurements of the organic, inorganic and total carbon yield of 190 Rock-Eval® thermal analysis to verify the ability of the Rock-Eval® thermal analysis to accurately measure the carbon amount of the samples.



To assess the effect of land cover on the Rock-Eval® parameters, we performed pairwise comparisons of medians by non-parametric Kruskal–Wallis tests ( $P < 0.05$ ) followed by Wilcoxon tests, with  $P < 0.05$  for each pair. The correction of p-values in the framework of the multiple comparisons was done with the Holm-Bonferroni method. Correlations between parameters were calculated using the Spearman method. We conducted a principal component analysis (using the R library FactoMineR) using all the observations and 11 pedoclimatic parameters: clay, total silt and total sand contents, pH in water, residual water content, carbonate content, mean annual temperature, mean annual precipitation, Tamm and Mehra-Jackson iron oxyhydroxides contents and C/N ratio (Fig. D). The data processing and statistical analysis were carried out with the R software (V4.1.2; R Core Team 2021. Packages integrated to R: base, datasets, graphics, grDevices, methods, stats, utils. Packages added: corrplot, car, ggplot2, ggpubr, factoextra, plot3D, rstatix, sf, tmap). The point maps of the Rock-Eval® Hydrogen Index and T50\_CO2\_PYR values were obtained using the tmap and sf R packages.

### 3 Results

#### 3.1 Carbon yields of Rock-Eval® thermal analysis

Figure 1a presents the TOCre6 plotted against TOCea. We observed a high correlation despite a few points far from the regression line ( $R^2=0.96$ ,  $n = 2037$ ) and an average carbon yield, corresponding to the slope of the regression, equal to 86%. Limiting the Rock-Eval® dataset to samples passing our quality check regarding the Rock-Eval® organic carbon yields (yields ranging from 0.7 and 1.3) left out 145 samples. Another sample was left out because of its TOC content: with a value of 0.57 g kg<sup>-1</sup>, this sample contains too little organic carbon for the data from the Rock-Eval® thermal analysis to be routinely exploitable (Khedim et al., 2021). A principal components analysis (PCA) conducted on all the topsoil samples showed no cluster for the samples with poor organic carbon yields (“rejected”) compared to the samples with good yields (“accepted”) (Fig. D). However, there was a significant difference between the means of the two groups for many pedoclimatic parameters. In particular, the total sand content was on average 76% higher in the rejected samples compared to accepted samples (101% higher for the coarse sand and 35% higher for the fine sand) and the carbonate content was also 67% higher in the rejected samples compared to accepted samples.

The remaining sample selection logically showed a better agreement between TOCre6 and TOCea, yet with on average lower TOCre6 values compared with TOCea (Rock-Eval® organic carbon yield of 0.87,  $R^2=0.99$ ,  $n = 1891$ ; Fig. 1b). The inorganic carbon content for this sample selection was slightly overestimated by the Rock-Eval® thermal analysis (Rock-Eval® inorganic carbon yield of 1.07,  $R^2=0.98$ ,  $n = 1891$ ). Finally, the total carbon content measured with Rock-Eval® thermal analysis for this sample selection is consistent with the total carbon measured using the elemental analysis (Rock-Eval® total carbon yield of 0.96,  $R^2=0.99$ ,  $n = 1891$ ).

**Figure 1: Carbon yields of Rock-Eval® thermal analysis. (a) Organic carbon yield: TOCre6 as a function of TOCea for all analyzed RMQS topsoil (0–30 cm) samples; (b) Organic carbon yield: TOCre6 as a function of TOCea for the RMQS topsoil samples limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3 ; (c) Inorganic carbon yield: MinC as a function of Cinorg for the RMQS topsoil samples limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3 ;**



225 and (d) Total carbon yield: TOCre6+Minc as a function of TOCea+Cinorg for the RMQS topsoil samples limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3.

### 3.2 Soil organic matter thermal stability in French topsoils and its relationships with land cover

The summary statistics of many different Rock-Eval® temperature parameters for the 1891 RMQS topsoil samples with satisfactory Rock-Eval® organic carbon yields are compiled in Supplementary Information (Table B).

230 Figure 2 shows the boxplots for the six selected parameters (T50\_HC\_PYR; T90\_HC\_PYR; T50\_CO2\_PYR; T50\_CO2\_OX; I-index; R-index) focusing on the four major land cover types.

235 **Figure 2: Effect of land cover on topsoil organic carbon thermal stability for the RMQS topsoil (0–30 cm) samples under the four major land covers in France: croplands, forests, grasslands, and vineyards & orchards. (a) T90\_HC\_PYR distribution; (b) T50\_CO2\_PYR distribution; (c) T50\_CO2\_OX distribution; (d) T50\_HC\_PYR distribution; (e) I-index distribution; (f) R-index distribution. Samples are limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3. For boxplots, the black midline of each box is the median. Lower and upper edges are respectively the 1<sup>st</sup> and 3<sup>rd</sup> quartiles. Lower and upper whiskers are respectively the maximum between the minimum value or the 1<sup>st</sup> quartile minus 1.5 times the interquartile range ( $\max[\min; Q1 - 1.5 \cdot (Q3 - Q1)]$ ) and the minimum between the maximum or the 3<sup>rd</sup> quartile plus 1.5 times the interquartile range ( $\min[\max; Q3 + 1.5 \cdot (Q3 - Q1)]$ ). Different letters indicate significant differences in the distribution of the values for the land uses with the Kruskal-Wallis test ( $P < 0.05$ ) and multiple Wilcoxon test ( $P < 0.05$ ).**

240 We observed similar results for the temperature parameters T90\_HC\_PYR, T50\_CO2\_PYR and T50\_CO2\_OX: thermal stability was significantly higher in croplands and vineyards & orchards compared to forests and grasslands. Topsoil organic carbon was slightly but significantly less thermally stable in forests than in grasslands (Fig. 2a, 2b, 2c). Notably, three other Rock-Eval® parameters related to SOC thermal stability in the HC\_PYR thermogram (T50\_HC\_PYR, I-index and R-index) showed a different response to land cover (Fig. 2d, 2e, 2f). The T50\_HC\_PYR and R-index indicated no significant difference in thermal stability in forests and croplands. The I-index indicated a value significantly lower in forests than in croplands.

### 3.3 Elemental stoichiometry of soil organic matter in French topsoils and its relationships with land cover

250 The summary statistics of different elemental stoichiometry parameters for the 1891 RMQS topsoil samples with satisfactory Rock-Eval® organic carbon yields are compiled in Supplementary Information (Table B). The HI (respectively OIre6 and C/N) mean value is 214 g HC.kg<sup>-1</sup> TOCre6 (respectively 177 g O<sub>2</sub>.kg<sup>-1</sup> TOCre6 and 12.05).

We observed significantly higher average values for the HI in both grasslands and forests compared to croplands and vineyards & orchards (Fig. 3, Fig. Cb). In contrast, grasslands and forests showed smaller values of OIre6 compared to croplands and vineyards & orchards (Fig. 3, Fig. Cc).

255 In addition, Figure 3 highlights that the distribution of the C/N ratio on the Rock-Eval® pseudo van Krevelen diagrams ( $HI=f(OIre6)$ ) depends on land cover. We observed a slight trend of the C/N ratio with the hydrogen and oxygen indices: the C/N ratio was higher for high HI and low OIre6. This trend was more pronounced for croplands and forests.

260 **Figure 3: Rock-Eval® pseudo van Krevelen diagrams ( $HI = f(OIre6)$ ) for the RMQS topsoil (0–30 cm) samples in (a) croplands; (b) grasslands; (c) forests; and (d) vineyards & orchards. Colours indicate the values of the C/N ratio. Samples are limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3.**



### 3.4 Correlations between Rock-Eval® indicators of SOM thermal stability and elemental stoichiometry and pedoclimate

Table 1 presents the Spearman correlation coefficient values of the Rock-Eval® temperature and stoichiometric parameters with the selected pedoclimatic variables. The three selected temperature parameters (T90\_HC\_PYR, T50\_CO2\_PYR, T50\_CO2\_OX) correlated significantly and positively with the clay content and negatively with the sand content. T90\_HC\_PYR and T50\_CO2\_PYR also correlated positively with silt content, however with smaller correlation coefficient values. They strongly and positively (correlation coefficient > 0.3) correlated with the water pH, the carbonate content and the cation exchange capacity, while the relationships with iron oxyhydroxides content were much lower. The three selected temperature parameters were all significantly positively correlated to Mean Annual Temperature (MAT) and negatively correlated to Mean Annual Precipitation (MAP), although the correlations were weak.

**Table 1: Spearman correlation coefficients of the Rock-Eval® temperature and stoichiometric parameters with pedoclimatic variables (TOCre6, particle-size distribution, pH in water, carbonate content, cation exchange capacity (CEC), iron oxyhydroxides, mean annual temperature (MAT) averaged over 1969-1999 and mean annual precipitation (MAP) averaged over 1969-1999) for the RMQS topsoil (0–30 cm) samples. The analysis was limited to samples with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3. Absolute values  $\geq 0.3$  are in bold. The asterisks indicate the p-value: 0 | \*\*\* | 0.001 | \*\* | 0.01 | \* | 0.05 | • | 0.1 | X.**

Regarding the indicators of SOM stoichiometry, HI and C/N correlated negatively with the clay and silt contents, contrary to OIre6 which correlated positively. HI and C/N also correlated negatively with the pH, the cation exchange capacity, and to a lesser extent with the carbonate content. They showed a slight negative correlation with the iron oxyhydroxides content measured by the Mehra-Jackson method. As for the thermal parameters, correlations with the climatic variables were on average smaller.

Additionally, the correlation coefficient of TOCre6 with HI was 0.35 (respectively -0.34 with OIre6, 0.37 with C/N, -0.26 with T90\_HC\_PYR, -0.21 with T50\_CO2\_PYR, -0.05 with T50\_CO2\_OX).

### 3.5 Distribution of some Rock-Eval® indicators of SOM thermal stability and elemental stoichiometry over the French mainland territory

Figure 4 shows the point maps of the HI and T50\_CO2\_PYR values over the French mainland territory. The missing topsoil samples (133 not included in the initial sample set and 146 rejected due to poor C yields) are distributed over the whole territory with some clusters in the north of the French Alps, north-east, Corsica, south-east and in the Landes. The first three clusters come from the 133 samples not included in the initial set. The Landes and south-east clusters are from both the absent samples and from the rejected samples: in particular, the soils in the Landes contain on average more sand, which is characteristic - as stated above - of the rejected samples. Visually, we noticed an autocorrelation of the values, HI and T50\_CO2\_PYR presenting on average opposite trends (the Spearman correlation coefficient between HI and T50\_CO2\_PYR is -0.69). Mountainous regions (notably the French Alps, the Pyrenees and the Massif Central) exhibit higher HI values and lower SOC thermal stability. Conversely, plain areas usually presented higher SOC thermal stability and lower HI values as in the Paris Basin and in the south-west and south-east part of the country. Brittany, Normandy and



the Landes are somewhat exceptions to this rule as they show high HI values and a relatively low SOC thermal stability. Figure 5 shows the land cover at each sampling site.

300 **Figure 4: Point maps of two Rock-Eval® parameters: (a) Hydrogen Index values and (b) T50\_CO2\_PYR values on the French mainland territory for the RMQS topsoil (0–30 cm) samples, along with (c) the map of the main regions used for the interpretation.**

**Figure 5: Map of the land cover at each sampling site. The number in brackets corresponds to the number of sites for each land cover in our final dataset (n=1891).**

## 4 Discussion

### 4.1 Carbon yields of Rock-Eval® thermal analysis

305 Our average organic carbon yield (0.86; Fig.1a) was in line with previous studies. Indeed, Disnar et al. (2003) obtained slightly higher yields (0.91), as well as Cécillon et al. (2018; 2021) (organic carbon yield from 0.90 to 0.96 depending on the sites) whereas Saenger et al. (2013) had lower yields (0.77). However, some samples presented high discrepancies between their TOC<sub>ea</sub> and TOC<sub>re6</sub> values. Samples with a TOC<sub>re6</sub> value strongly differing from its corresponding TOC<sub>ea</sub> value were  
310 organically re-analyzed using Rock-Eval®, which confirmed their first TOC<sub>re6</sub> measurement. The outliers regarding the organic carbon yield were thus not related to a problem in their Rock-Eval® measurement. These very different values, which concern a few dozens of samples, could have different origins such as error on sample labeling, aliquoting, grinding or storage conditions. Indeed, for the same sample, the powders used for the elemental analysis and the Rock-Eval® thermal analysis did not come from the same aliquot. In addition, the elemental analyses were performed shortly after sampling, whereas the samples analyzed in Rock-Eval® were stored for about fifteen years. We can therefore expect slightly better  
315 yields when elementary and Rock-Eval® analysis are performed with less time between both, and on the exact same powders. This is what we plan for the samples of the second RMQS sampling campaign. The very different values between TOC<sub>ea</sub> and TOC<sub>re6</sub> could also be due, for some samples, to a mismeasurement of the total carbonate content, leading to a miscalculation of the inorganic and organic carbon contents. This hypothesis could be plausible, as the mean value of the carbonate content was significantly higher in the rejected samples. The last hypothesis originates from the high content of  
320 sand in the rejected samples: sandy samples are more heterogeneous, thus the material used to determine the TOC<sub>ea</sub> is more likely to differ from the one used to determine the TOC<sub>re6</sub>, than when sand content is lower. Moreover, the physical state of organic matter in sandy soils can be different from other soils. Disnar et al. (2003) encountered “pellets” of SOM in sandy soils, which can strongly influence the results of TOC<sub>ea</sub> and TOC<sub>re6</sub>.

325 The samples presenting a high discrepancy between TOC<sub>ea</sub> and TOC<sub>re6</sub> were not considered further in the analysis. As stated above, we restricted our study to the samples with organic carbon yield ranging from 0.7 to 1.3. This subjective threshold is a quality threshold to ensure that the samples analyzed using Rock-Eval® were the same as the samples analyzed using elemental analysis on which rely all studies conducted on the first campaign of the RMQS. This selection only marginally improved the average organic carbon yield (0.87; Fig. 1b) and organic carbon was still underestimated by



Rock-Eval®. Conversely, inorganic carbon yield was slightly overestimated (1.07; Fig. 1c). As a result, the yield of total  
330 carbon (organic + inorganic carbon) was close to 1.00 (Fig. 1d). This suggests that almost all sample carbon is detected by  
the Rock-Eval® machine in the five thermograms but that a small part of the organic carbon is erroneously attributed to  
inorganic carbon. This may be due to a slight misplacement of the boundary between organic and inorganic carbon, probably  
in the S3 and S3CO signals. Also, the S3'CO signal is attributed half to organic carbon and half to inorganic carbon due to  
335 of the Rock-Eval® peaks). Of note, as MinC and TOCre6 are very well correlated to Cinorg and TOCea ( $R^2 > 0.98$ ), it  
should therefore be possible to draw a correction formula to assess TOCea and Cinorg using Rock-Eval® with high  
accuracy. This would allow determining simultaneously in less than 1 h organic C and inorganic C with no risk of error due  
to erroneous decarbonation.

#### 4.2 Thermal stability of soil organic carbon in French topsoils

340 We have observed that the thermal stability defined according to different Rock-Eval® parameters varies in French topsoils.  
We can investigate whether these variations are consistent with our knowledge of SOC biogeochemical stability. SOC  
biogeochemical stability is on average higher in croplands and vineyards compared to forest or grassland soils (Poeplau and  
Don, 2013). Indeed, fresh organic carbon inputs to soil are usually higher in forest and grassland compared to croplands  
where human exportation of biomass is higher (Murty et al., 2002). As a result, SOC fractions with lower mean residence  
345 time in soils and lower thermal stability can be more abundant in forests and grasslands compared to croplands. For instance,  
several studies reported that carbon in particulate organic matter (a relatively more labile form of SOC) contributes more to  
total SOC in forest and grassland compared to cropland (e.g. Guo and Gifford, 2002; Poeplau et al., 2011; Poeplau and Don,  
2013; Lugato et al. 2021). Moreover, agricultural practices may also speed up SOC mineralization further limiting the  
accumulation of labile SOC fractions. For instance, Balesdent et al. (1990) observed that the tillage practices lead to a  
350 significantly higher mineralization than no tillage. Combining the effects of lower carbon inputs and mineralization-  
enhancing practices, croplands contain on average less biogeochemically labile SOC than forests and grasslands.

Thermal stability, as assessed using T90\_HC\_PYR, T50\_CO2\_PYR and T50\_CO2\_OX, was the highest in vineyards,  
orchards and croplands compared to forest and grassland soils (Fig. 2). These results suggest that, overall, SOC thermal  
stability as assessed using these Rock-Eval® parameters is related to SOC biogeochemical stability. This is in good  
355 agreement with previous results obtained on smaller datasets (Barré et al., 2016; Poeplau et al., 2019; Cécillon et al., 2021).  
On the contrary, there was no consistent relationship between thermal stability and expected biogeochemical stability when  
the thermal stability was measured using T50\_HC\_PYR, R-index and I-index (Fig. 2). Cécillon et al. (2021) reported for  
soils with highly contrasted biogeochemical stability that the relationship between thermal stability and biogeochemical  
stability was weaker for T50\_HC\_PYR, R-index and I-index. Our results showed that this relationship even disappears when  
360 considering data sets with more heterogeneous topsoil samples. The use of the Rock-Eval® temperature parameters



T90\_HC\_PYR, T50\_CO2\_PYR and T50\_CO2\_OX should therefore be preferred when seeking to measure thermal stability indicators directly related to biogeochemical stability.

T90\_HC\_PYR, T50\_CO2\_PYR and T50\_CO2\_OX were all strongly and positively correlated to clay content and negatively correlated to sand content (Table 1). In a previous study, Soucémariadin et al. (2018) did not observe any correlation  
365 between T50\_CO2\_OX and clay or sand content, however, their study was conducted on forest soils only and on a much reduced number of study sites. Soil clay fractions interact with microbial compounds which results in the formation of organo-mineral complexes which SOC has a high biogeochemical stability (e.g. Lehmann and Kleber, 2015). We can therefore hypothesize that clay-rich soils are also richer in biogeochemically stable carbon. The positive correlation between clay content and SOC thermal stability, and the good correlations between CEC, which depends on the first order of the clay  
370 content, and SOC thermal stability would then be another illustration of the link between SOC thermal and biogeochemical stabilities. Iron oxides are mineral compounds that are also supposed to protect SOC from decomposition. To this respect, the inconsistent (Mehra-Jackson Iron) or even negative correlations (Tamm Iron) between T90\_HC\_PYR, T50\_CO2\_PYR and T50\_CO2\_OX and iron oxides were not expected. These weak correlations could be attributed to the fact that the range of iron oxides contents is relatively small in our set of topsoils.

375 T90\_HC\_PYR, T50\_CO2\_PYR and T50\_CO2\_OX were all positively correlated to pH. Such a correlation between T50\_CO2\_OX and pH was already observed by Soucémariadin et al. (2018) for a set of French forest soils. Acidity may protect SOM from degradation by microorganisms (Clivot et al., 2021), by reducing their activity, which is actually observed in low pH acidic bogs. We can therefore hypothesize that acidity slows down SOM mineralization which can favor the accumulation of labile SOC components. As these labile SOC fractions would appear as thermally unstable, it would explain  
380 the positive relationship between pH and Rock-Eval® indicators of SOC thermal stability.

T90\_HC\_PYR, T50\_CO2\_PYR and T50\_CO2\_OX showed weak but significant positive correlations with MAT averaged over 1969-1999 (Fig. 5). Such a correlation has also been observed in Soucémariadin et al. (2018) for French forest soils. As soil microbial activity and thus SOC mineralization increase with temperature (Rey and Jarvis, 2006), we can expect SOC labile fractions to be more rapidly processed at higher temperature. It would be in line with the observed positive  
385 correlations between MAT and the three selected thermal stability indicators. The relatively weak (Spearman rho value below 0.2) correlations can be due to the fact that MAT will also play on carbon inputs to the soil. Indeed, if higher SOC mineralization was balanced by increased biomass inputs it would mess up the relation between MAT and SOC biogeochemical stability. In a similar way, the weak negative correlation between MAP and thermal stability may be explained by the complex effect of MAP on SOC biogeochemical stability: increased soil moisture stimulates SOC  
390 processing up to a certain point (Moyano et al., 2013) and influences net primary production and therefore the soil carbon inputs. In any case, the relationships between SOC and MAP or MAT are hard to disentangle (Chen et al., 2019). Another explanation for the weak values is that the climatic data was obtained on an 8km × 8km grid and do not have the same precision as if a weather station had been deployed at each site. This probably adds noise to the correlation.



The point map representing SOC thermal stability over mainland France (Fig. 4b) illustrates the relationships between SOC  
395 thermal stability, land cover, climate and pedological variables. Mountainous regions (e.g. Massif Central, Alps, Pyrenees)  
where forest, grassland, and low MAT dominate, and presented by Martin et al. (2011) as with relatively high SOC contents,  
had a lower SOC thermal stability. Plains dominated by croplands with intensive agricultural practices and with relatively  
low SOC contents such as the Paris Basin showed high SOC thermal stability. The southern part of France with warmer  
MAT, dominated by vineyards and croplands, and relatively low SOC contents also presented high SOC thermal stability.  
400 The lower SOC thermal stabilities observed in Brittany and Normandy (which are agricultural regions) could be explained  
by the higher proportion of livestock. Therefore, in addition to the presence of grasslands in these regions, the cultivated  
soils in Brittany and Normandy are more likely to receive repeated application of exogenous organic matter.

#### 4.3 Elemental stoichiometry of soil organic matter in French topsoils

Higher values of HI and lower values of OIre6 were observed in forests and grasslands compared to croplands and  
405 vineyards. This trend was observed in previous studies (Disnar et al., 2003; Saenger et al., 2013; Sebag et al., 2016). It also  
confirms that HI and OIre6 can be good proxies of SOC biogeochemical stability. Indeed, as previously observed,  
biogeochemically stable SOC is more oxidized and H-depleted (Barré et al., 2016; Poeplau et al., 2019; Cécillon et al.,  
2021).

The pseudo van Krevelen diagrams (Fig. 3) showed a high variability of the C/N ratio between land cover classes: the C/N  
410 ratio was higher in forest topsoils than in grasslands, as well as in croplands and vineyards. This is classically explained by  
the fact that SOC is on average less processed in forests and grasslands compared to croplands and vineyards (Cotrufo et al.,  
2019), as well as by the higher C/N ratio of the biomass inputs to soil in forests. Indeed, the biotransformation of organic  
matter tends to lower its C/N ratio and oxidize it (Cleveland and Liptzin, 2007). This is in good agreement with the observed  
trends of decreasing HI and increasing OIre6 with decreasing C/N (Fig. 3).

SOM elemental stoichiometry presented correlation patterns with land cover, climate and pedological variables that were  
415 similar to those observed for SOM thermal stability. HI and OIre6 are respectively negatively and positively correlated to pH  
(Table 1) as previously observed by Soucémarianadin et al. (2018) in French forest soils. This would be in line with acidity  
slowing down the mineralization of H-enriched labile SOC fractions (Clivot et al., 2019). The negative correlation between  
clay content and HI could be explained by the fact the presence of clays can promote the protection of microbially processed  
420 H-depleted SOM. Similarly to what was observed for SOM thermal stability, relationships between elemental stoichiometry  
and climate variables are weak, probably because climate plays on both soil carbon inputs and outputs in opposite ways  
(climate conditions enhancing SOC mineralization usually also enhance fresh SOM inputs).

The point map of HI in mainland France (Fig. 4a) illustrated the effect of land cover, climate and pedological variables on  
SOM elemental stoichiometry. Regions dominated by grassland and forest (Fig. 5) such as mountainous regions, the Landes  
425 forest or the forest-dominated east part of France were characterized by a relatively H-enriched SOM. Conversely, regions  
dominated by croplands, vineyards & orchards, and by high MAT were characterized by a relatively H-depleted SOM. Both



point maps of thermal stability and HI (Fig. 4) also illustrated the relationships previously observed between these Rock-Eval® parameters (Barré et al., 2016; Cécillon et al., 2021).

## 5 Conclusion

430 This study is an unprecedented effort to make widespread thermal analysis measurements on a national soil quality monitoring network. It demonstrated that Rock-Eval® may be used as a rapid and cost-effective method to assess the thermal stability and elemental stoichiometry of SOM on national soil monitoring networks. Our results highlighted the influence of land cover and pedoclimatic variables on SOM thermal stability and elemental stoichiometry. They suggested that some Rock-Eval® temperature parameters describing SOC thermal stability (T90\_HC\_PYR, T50\_CO2\_PYR and  
435 T50\_CO2\_OX) could be used as reliable proxies of SOC biogeochemical stability, while others parameters could not (T50\_HC\_PYR, R-index and I-index). Our study also opened wide perspectives for future research. In the short term, these Rock-Eval® results on French topsoils can be used as input to the PARTYSOC machine-learning model (Cécillon et al. 2021) to infer the size of the centennially stable SOC fraction. They can also be compared to other proxies of SOC biogeochemical stability such as SOM physical fractionation results. In the medium term, it will be interesting to test  
440 whether this analytical information can be used to improve the accuracy of SOC stock evolution simulations at the scale of a national soil monitoring network, as it was observed for the AMG model of SOC dynamics in several French long term agronomic experiments (Kanari et al., 2022).

## Data availability

Data on basic soil properties are freely available from the GIS Sol dataverse website:  
445 <https://data.inrae.fr/dataset.xhtml?persistentId=doi:10.15454/BNCXYB>.

## Author contribution

Amicie Delahaie, Lauric Cécillon and Pierre Barré gathered the sample collection and ensured that samples were properly ground. Florence Savignac and François Baudin produced the Rock-Eval® thermal analyses. Dominique Arrouays, Antonio Bispo, Line Boulonne, Claudy Jolivet, Manuel Martin, Céline Ratié and Nicolas Saby provided the detailed pedoclimatic  
450 data. Nicolas Saby produced the point maps. Amicie Delahaie processed and interpreted the data with the contribution of all co-authors. Amicie Delahaie, Pierre Barré and Lauric Cécillon wrote the manuscript with contribution of all the co-authors.

## Competing interests

The authors declare that they have no conflict of interest.



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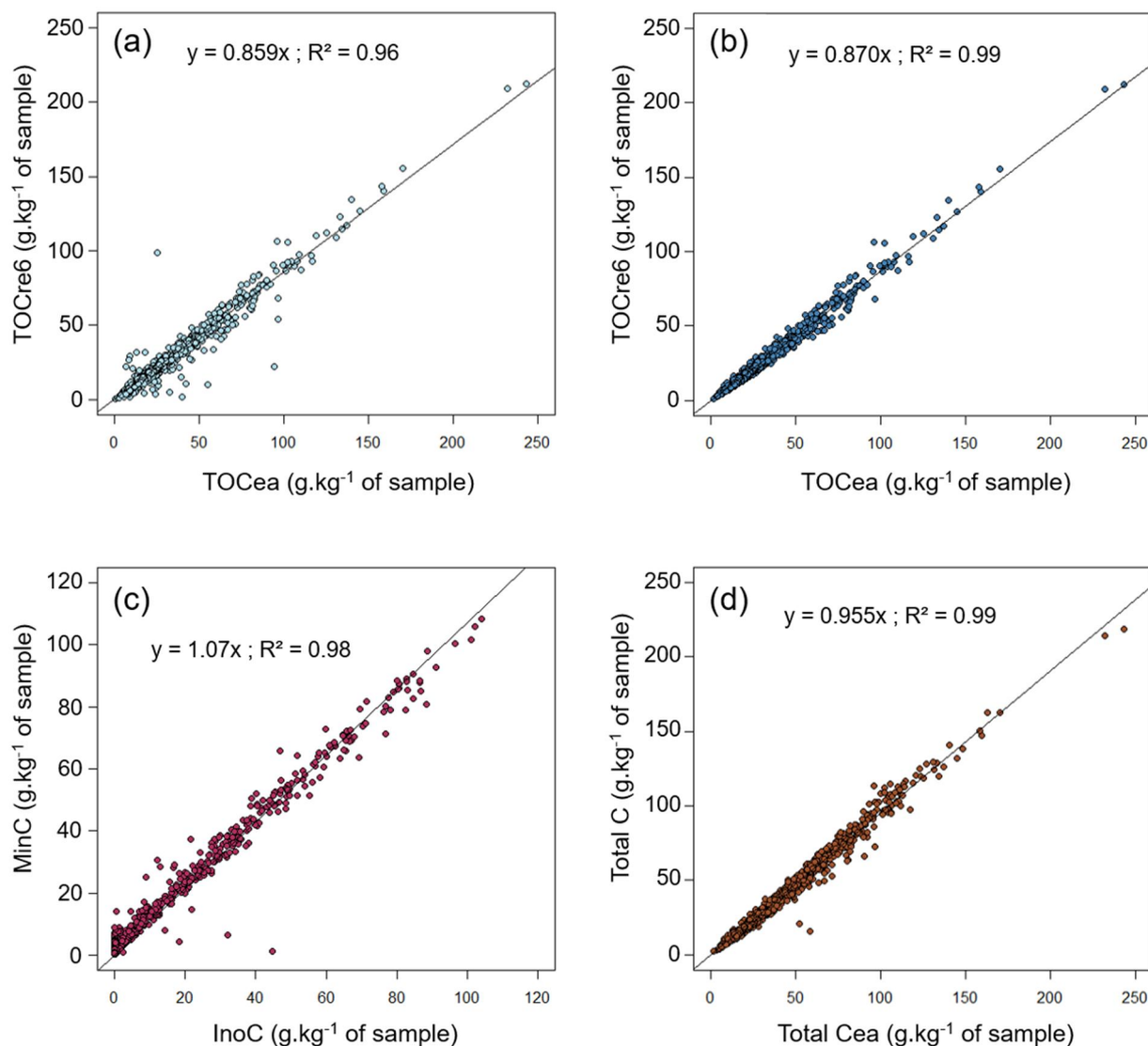
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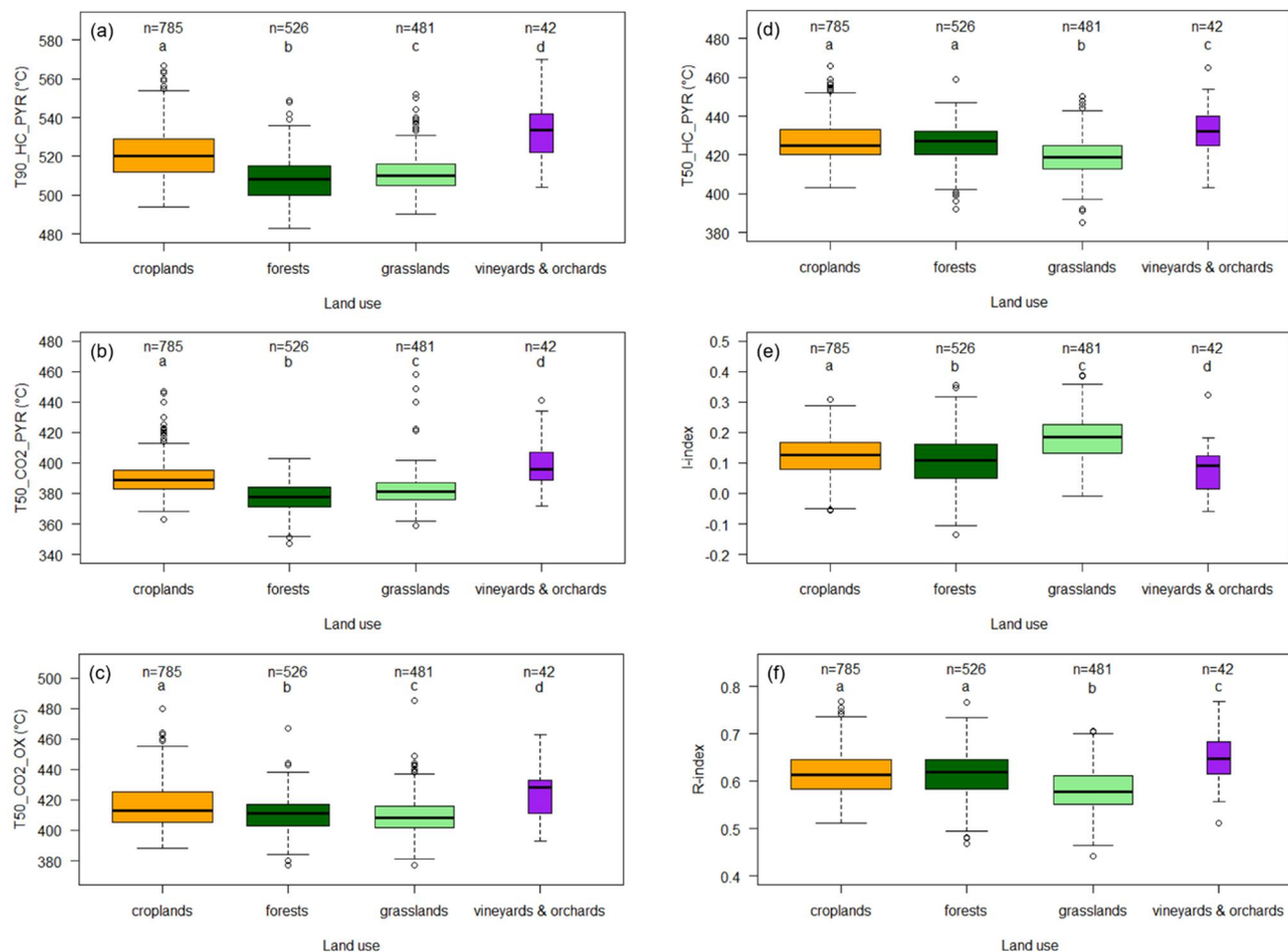
	T90_HC_PYR	T50_CO2_PYR	T50_CO2_OX	HI	Olre6	C/N
<b>TOCre6</b> (n=1891)	-0.26 ***	-0.21 ***	-0.05 *	<b>0.35</b> ***	<b>-0.34</b> ***	<b>0.37</b> ***
<b>clay</b> (n=1891)	<b>0.46</b> ***	<b>0.56</b> ***	<b>0.45</b> ***	<b>-0.35</b> ***	<b>0.33</b> ***	-0.27 ***
<b>silt (total)</b> <b>(fine coarse)</b> (n=1891)	0.13 *** 0.12   0.09 ***   ***	0.20 *** 0.29   0.12 ***   ***	-0.04 • 0.03   -0.09 x   ***	-0.18 *** -0.16   -0.17 ***   ***	0.26 *** 0.23   0.23 ***   ***	<b>-0.31</b> *** -0.23   -0.28 ***   ***
<b>sand (total)</b> <b>(fine coarse)</b> (n=1891)	<b>-0.36</b> *** -0.17   -0.34 ***   ***	<b>-0.42</b> ***	-0.25 ***	<b>0.31</b> ***	<b>-0.34</b> ***	<b>0.35</b> ***
<b>water pH</b> (n=1891)	<b>0.71</b> ***	<b>0.73</b> ***	<b>0.44</b> ***	<b>-0.42</b> ***	<b>0.39</b> ***	<b>-0.52</b> ***
<b>carbonates</b> (n=1891)	<b>0.53</b> ***	<b>0.56</b> ***	<b>0.45</b> ***	-0.20 ***	0.24 ***	-0.28 ***
<b>CEC</b> (n=1891)	<b>0.60</b> ***	<b>0.56</b> ***	<b>0.46</b> ***	<b>-0.36</b> ***	<b>0.32</b> ***	<b>-0.36</b> ***
<b>Free iron (Tamm)</b> (n=1622)	-0.16 ***	-0.13 ***	-0.26 ***	-0.06 *	0.09 ***	-0.07 **
<b>Free iron (Mehra-Jackson)</b> (n=1621)	0.08 **	<b>0.36</b> ***	-0.05 *	<b>-0.33</b> ***	<b>0.35</b> ***	-0.11 ***
<b>MAT (1969-1999)</b> (n=1891)	0.12 ***	0.24 ***	0.10 ***	-0.20 ***	0.06 **	-0.11 ***
<b>MAP (1969-1999)</b> (n=1891)	-0.25 ***	-0.09 ***	-0.20 ***	0.14 ***	-0.10 ***	0.21 ***

650 Table 1: Spearman correlation coefficients of the Rock-Eval® temperature and stoichiometric parameters with pedoclimatic  
 variables (TOCre6, particle-size distribution, pH in water, carbonate content, cation exchange capacity (CEC), iron  
 oxyhydroxides, mean annual temperature (MAT) averaged over 1969-1999 and mean annual precipitation (MAP) averaged over  
 1969-1999) for the RMQS topsoil (0–30 cm) samples. The analysis was limited to samples with Rock-Eval® organic carbon yields  
 ranging from 0.7 to 1.3. Absolute values  $\geq 0.3$  are in bold. The asterisks indicate the p-value: 0 | \*\*\* | 0.001 | \*\* | 0.01 | \* | 0.05 | • |  
 655 0.1 | X.

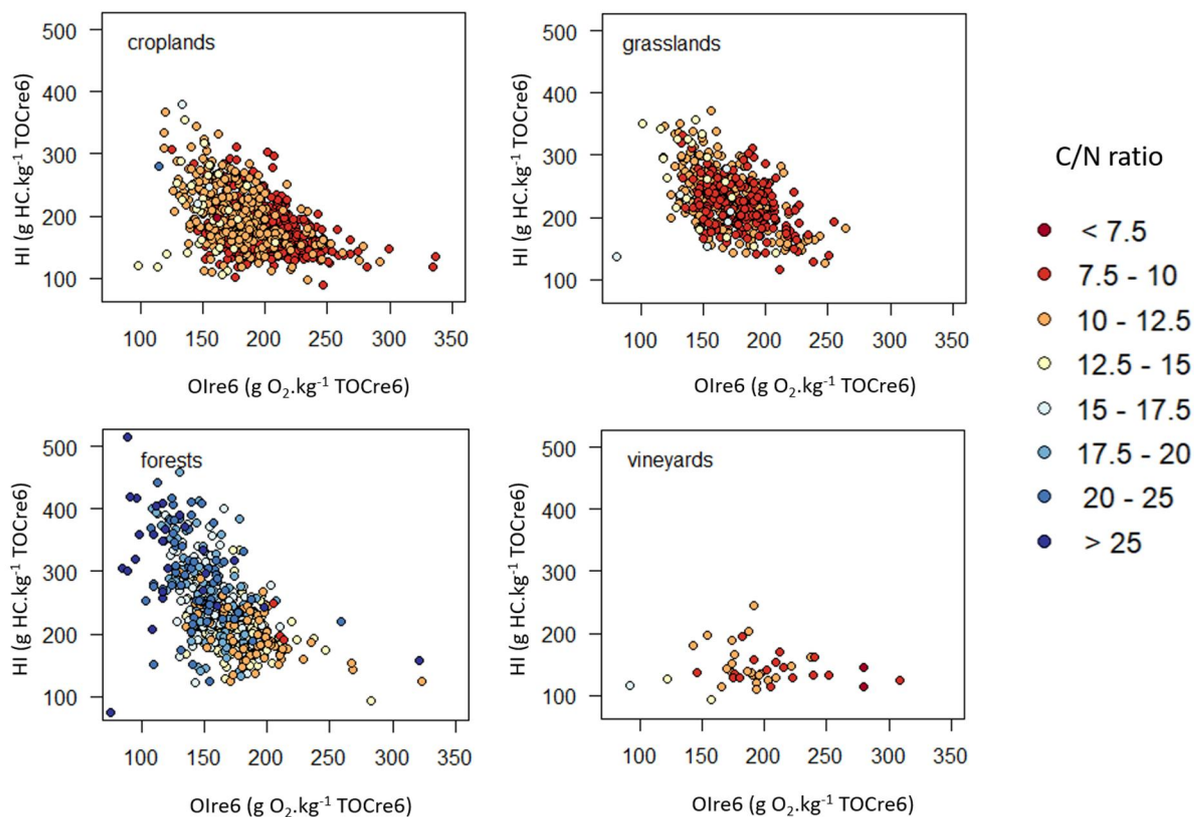


660 **Figure 1: Carbon yields of Rock-Eval® thermal analysis. (a) Organic carbon yield: TOCre6 as a function of TOCea for all analyzed RMQS topsoil (0–30 cm) samples; (b) Organic carbon yield: TOCre6 as a function of TOCea for the RMQS topsoil samples limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3 ; (c) Inorganic carbon yield: MinC as a function of Cinorg for the RMQS topsoil samples limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3 ; and (d) Total carbon yield: TOCre6+Minc as a function of TOCea+Cinorg for the RMQS topsoil samples limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3.**

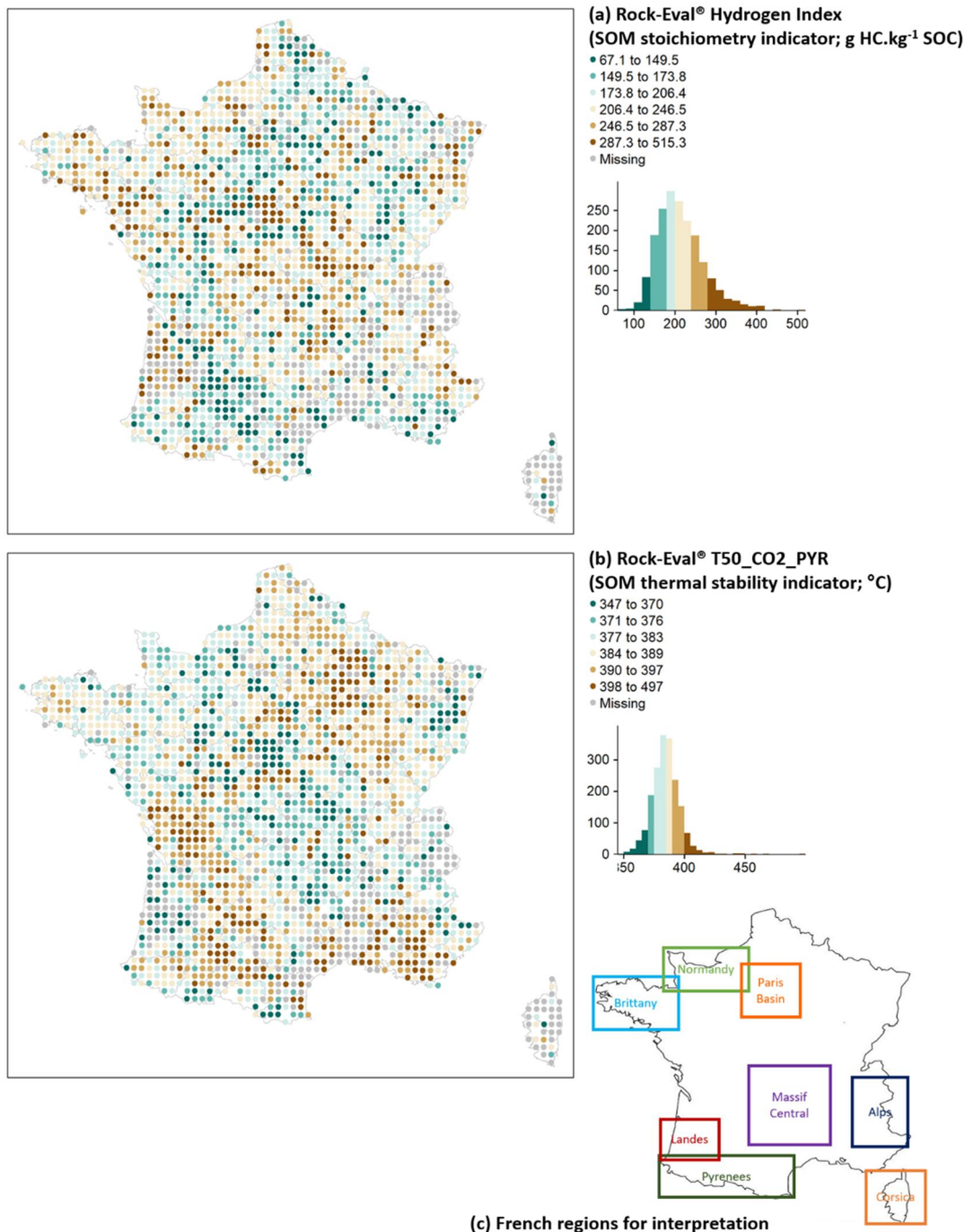
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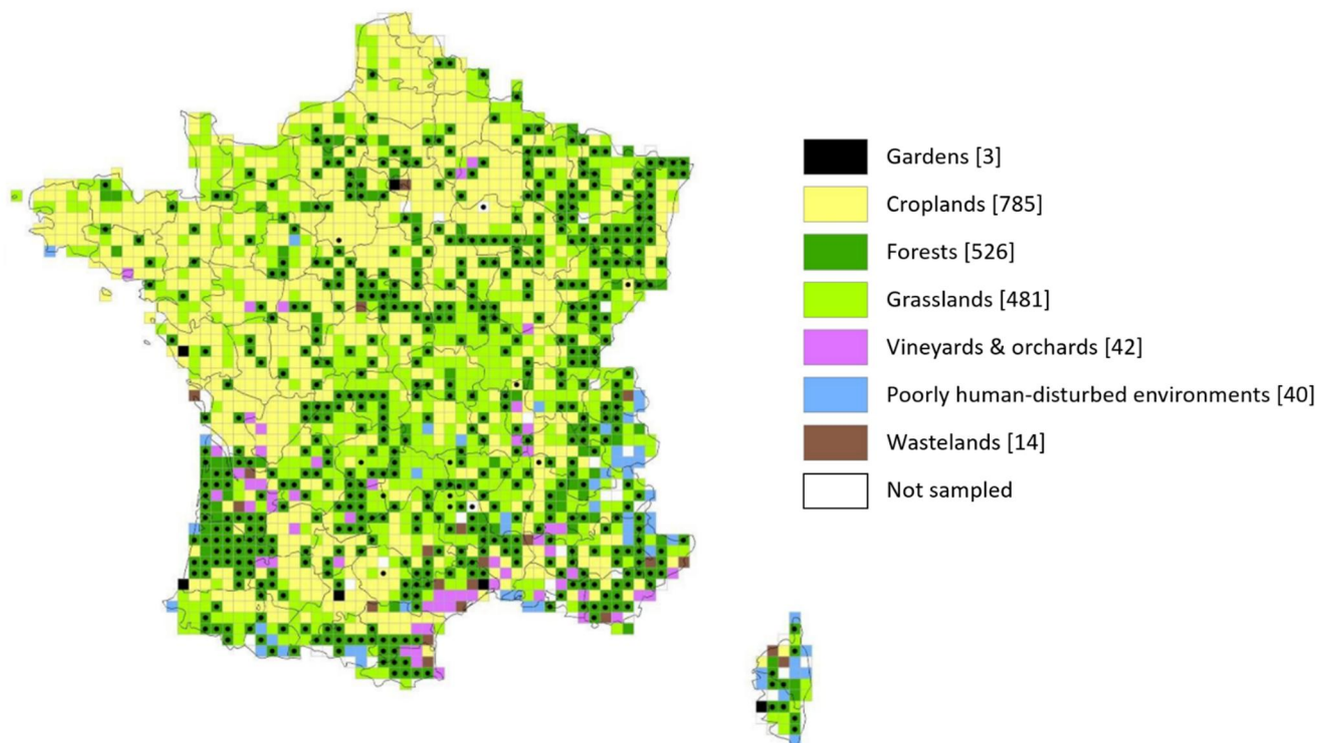
670 **Figure 2: Effect of land cover on topsoil organic carbon thermal stability for the RMQS topsoil (0–30 cm) samples under the four major land covers in France: croplands, forests, grasslands, and vineyards & orchards. (a) T90\_HC\_PYR distribution; (b)**  
 675 **T50\_CO2\_PYR distribution; (c) T50\_CO2\_OX distribution; (d) T50\_HC\_PYR distribution; (e) I-index distribution; (f) R-index distribution. Samples are limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3. For boxplots, the black midline of each box is the median. Lower and upper edges are respectively the 1<sup>st</sup> and 3<sup>rd</sup> quartiles. Lower and upper whiskers are respectively the maximum between the minimum value or the 1<sup>st</sup> quartile minus 1.5 times the interquartile range (max[ $\min$ ;  $Q1 - 1.5 \cdot (Q3 - Q1)$ ]) and the minimum between the maximum or the 3<sup>rd</sup> quartile plus 1.5 times the interquartile range (min[ $\max$ ;  $Q3 + 1.5 \cdot (Q3 - Q1)$ ]). Different letters indicate significant differences in the distribution of the values for the land uses with the Kruskal-Wallis test ( $P < 0.05$ ) and multiple Wilcoxon test ( $P < 0.05$ ).**



680 **Figure 3: Rock-Eval® pseudo van Krevelen diagrams ( $HI = f(Olre6)$ ) for the RMQS topsoil (0–30 cm) samples in (a) croplands; (b) grasslands; (c) forests; and (d) vineyards & orchards. Colours indicate the values of the C/N ratio. Samples are limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3.**



**Figure 4: Point maps of two Rock-Eval® parameters: (a) Hydrogen Index values and (b) T50\_CO2\_PYR values on the French mainland territory for the RMQS topsoil (0–30 cm) samples, along with (c) the map of the main regions used for interpretation.**



**Figure 5: Map of the land cover at each sampling site (Jolivet, 2011). The number in brackets corresponds to the number of sites for each land cover in our final dataset (n=1891).**



## 690 Appendices

parameter	unit	formula	description
TOCre6	g.kg <sup>-1</sup> sample	$PC \times 10 + S4CO \times \frac{12}{28}$	total organic carbon
MinC	g.kg <sup>-1</sup> sample	$S3' \times \frac{12}{44} + \frac{S3'CO}{2} \times \frac{12}{28} + S5 \times \frac{12}{44}$	total inorganic carbon
PC	g.kg <sup>-1</sup> sample	$\frac{(S1 + S2) \times 0.83 + S3 \times \frac{12}{44} + (S3CO + \frac{S3'CO}{2}) \times \frac{12}{28}}{10}$	amount of pyrolyzable organic carbon
PC/TOCre6	no unit	$\frac{PC}{TOCre6}$	ratio of pyrolyzable organic carbon over total organic carbon
PseudoS1	g.kg <sup>-1</sup> sample	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions)	carbon released during the first pyrolysis isotherm
S2	g.kg <sup>-1</sup> sample	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions)	carbon released as hydrocarbons during the pyrolysis except during the first isotherm
S2/PC	no unit	$\frac{S2}{PC}$	ratio of carbon released as hydrocarbons during the pyrolysis except during the first isotherm over the pyrolyzable organic carbon
T50_HC_PYR	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 50 % of the hydrocarbon effluents have been emitted during the pyrolysis ramp (the initial isotherm is excluded; the integration ends at 650°C)
T70_HC_PYR	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 70 % of the hydrocarbon effluents have been emitted during the pyrolysis ramp (the initial isotherm is excluded; the integration ends at 650°C)
T90_HC_PYR	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 90 % of the hydrocarbon effluents have been emitted during the pyrolysis ramp (the initial isotherm is excluded; the integration ends at 650°C)
T30_CO2_PYR	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 30 % of the CO2 have been emitted during the pyrolysis ramp (the beginning isotherm is excluded; the integration ends at 560°C)
T50_CO2_PYR	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 50 % of the CO2 have been emitted during the pyrolysis ramp (the beginning isotherm is excluded; the integration ends at 560°C)
T70_CO2_PYR	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 70 % of the CO2 have been emitted during the pyrolysis ramp (the beginning isotherm is excluded; the integration ends at 560°C)
T90_CO2_PYR	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 90 % of the CO2 have been emitted during the pyrolysis ramp (the beginning isotherm is excluded; the integration ends at 560°C)
T50_CO_PYR	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 50 % of the CO have been emitted during the pyrolysis ramp (the beginning isotherm is excluded; the integration ends at 560°C)
T50_CO2_OX	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 50 % of the CO2 have been emitted during the oxidation phase (the integration ends at 611°C)
T70_CO2_OX	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 70 % of the CO2 have been emitted during the oxidation phase (the integration ends at 611°C)
T90_CO2_OX	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 90 % of the CO2 have been emitted during the oxidation phase (the integration ends at 611°C)
T50_CO_OX	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 50 % of the CO have been emitted during the oxidation phase (the integration ends at 850°C)
T70_CO_OX	°C	Integration of the thermogram (see Behar et al., 2001 for thermograms descriptions) to obtain temperature	temperature at which 70 % of the CO have been emitted during the oxidation phase (the integration ends at 850°C)
I-index	no unit	Integration of the thermogram (see Sebag et al., 2016 for boundaries)	related to the thermolabile organic carbon released as hydrocarbon effluents, see Sebag et al., 2016
R-index	no unit	Integration of the thermogram (see Sebag et al., 2016 for boundaries)	proportion of thermostable organic carbon released as hydrocarbon effluents after 400°C, see Sebag et al., 2016
HI	g HC.kg <sup>-1</sup> TOCre6	$\frac{S2 \times 100}{TOCre6}$	ratio of emitted hydrocarbons to TOCre6
OIre6	g O <sub>2</sub> .kg <sup>-1</sup> TOCre6	$\frac{16}{28} \times \frac{S3CO \times 100}{TOCre6} + \frac{32}{44} \times \frac{S3 \times 100}{TOCre6}$	ratio of organic oxygen to TOCre6

**Table A: Description of the Rock-Eval® parameters and their calculation.**

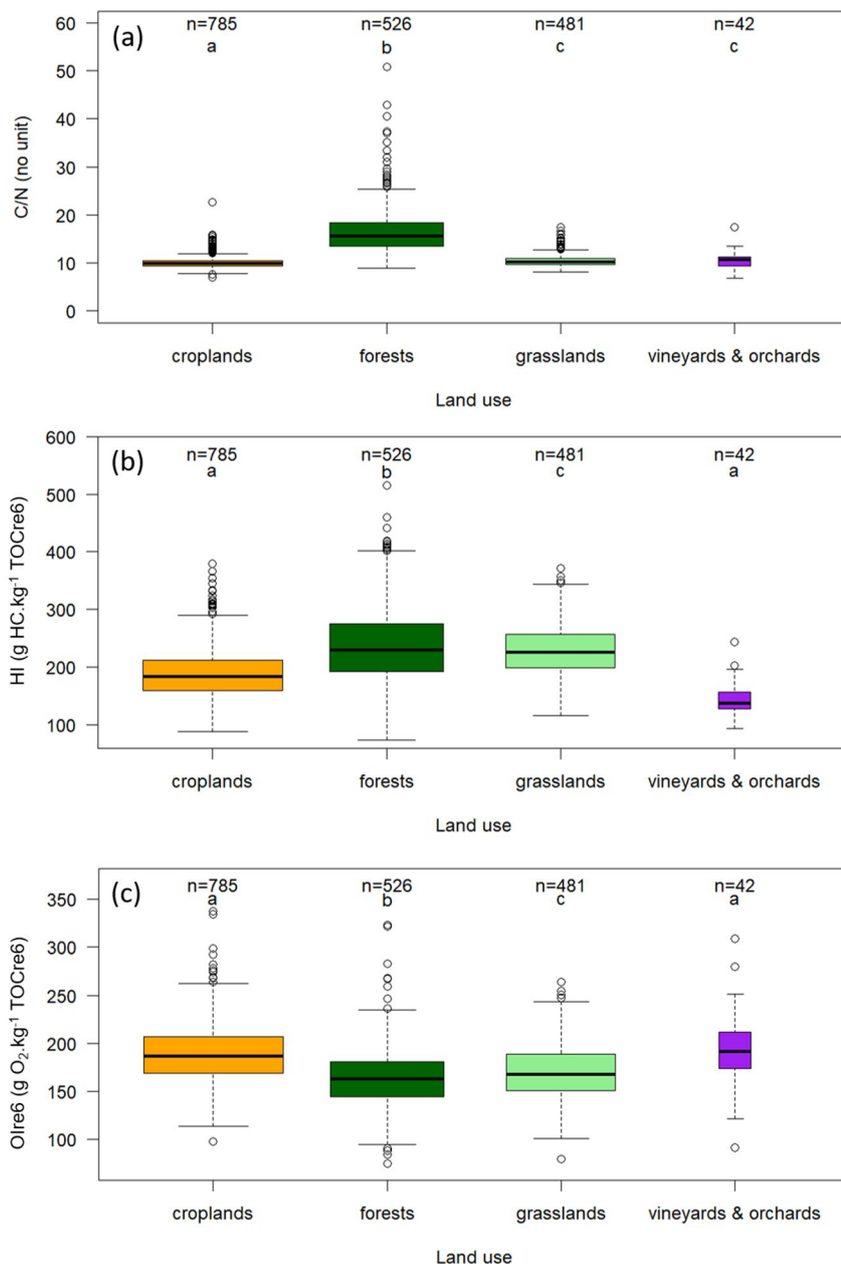


	min					mean					max				
	total	croplands	grasslands	forests	vineyards & orchards	total	croplands	grasslands	forests	vineyards & orchards	total	croplands	grasslands	forests	vineyards & orchards
C_N	6,9	7,0	8,2	8,9	6,9	12,1	10,1	10,5	16,7	10,5	50,8	22,6	17,4	50,8	17,4
T50_HC_PYR	385	403	385	392	403	424	426	419	425	432	466	466	450	459	465
T70_HC_PYR	438	445	438	438	453	464	466	461	463	473	508	508	488	492	506
T90_HC_PYR	483	494	490	483	504	515	521	511	508	534	572	567	552	549	570
T30_CO2_PYR	311	325	318	311	327	337	341	335	331	348	448	381	385	350	379
T50_CO2_PYR	347	363	359	347	372	384	390	382	377	399	497	447	458	403	441
T70_CO2_PYR	389	411	408	389	423	437	442	435	429	453	527	500	509	461	496
T90_CO2_PYR	462	485	484	462	495	505	509	505	499	517	550	542	544	529	542
T50_CO_PYR	199	199	338	377	386	404	402	401	408	404	472	472	470	464	427
T50_CO_OX	326	350	326	329	370	401	410	393	391	433	586	586	529	492	545
T70_CO_OX	356	386	363	356	408	457	473	448	439	504	690	666	690	551	611
T50_CO2_OX	377	388	377	377	393	413	416	409	410	425	493	480	485	467	463
T70_CO2_OX	404	417	406	404	436	461	468	455	454	484	554	539	534	512	528
T90_CO2_OX	444	465	464	444	518	530	537	528	521	548	597	585	577	563	577
PseudoS1	0,02	0,02	0,03	0,04	0,02	0,14	0,09	0,15	0,18	0,08	0,98	0,33	0,64	0,98	0,22
S2	0,14	0,59	0,85	0,14	0,34	4,22	2,30	5,12	5,95	1,18	61,6	8,26	27,3	32,9	5,53
S2_PC	0,31	0,31	0,46	0,38	0,40	0,65	0,62	0,68	0,69	0,54	0,9	0,82	0,84	0,9	0,68
HI	67	88	116	74	94	214	190	229	240	144	515	379	372	515	244
Olre6	75	98	80	75	92	177	189	171	163	198	337	337	264	323	325
HI_Olre6	0,32	0,36	0,51	0,33	0,39	1,28	1,05	1,39	1,57	0,76	5,82	3,06	3,46	5,82	1,27
PC	0,33	0,95	1,56	0,33	0,77	6,14	3,62	7,32	8,45	2,12	73,5	11,9	36,3	43,6	8,09
PC_TOCre6	0,1	0,14	0,16	0,1	0,15	0,27	0,25	0,28	0,28	0,22	0,48	0,4	0,38	0,48	0,3
TOCre6	1,1	3,6	5,4	1,1	2,8	22,5	14,1	26,3	30,2	9,5	213	46,5	127	144	27,3
MINC	0,2	0,2	0,3	0,2	0,5	8,0	9,7	6,4	6,4	12,4	108	108	85,8	97,9	58,3
R_index	0,44	0,51	0,44	0,47	0,51	0,61	0,62	0,58	0,61	0,65	0,77	0,77	0,71	0,77	0,77
I_index	-0,14	-0,06	-0,01	-0,14	-0,06	0,13	0,12	0,18	0,11	0,08	0,39	0,31	0,39	0,36	0,32

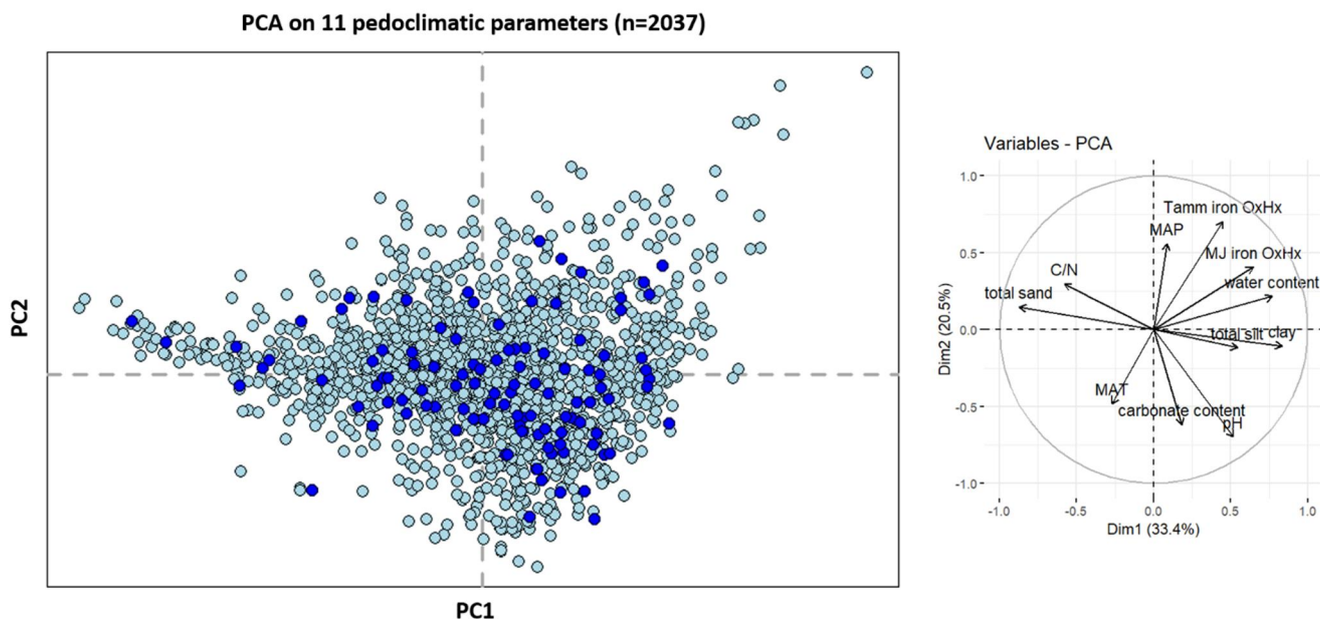


	First quartile					Median					Third quartile					Standard deviation				
	total	croplands	grasslands	forests	vineyards & orchards	total	croplands	grasslands	forests	vineyards & orchards	total	croplands	grasslands	forests	vineyards & orchards	total	croplands	grasslands	forests	vineyards & orchards
C_N	9,7	9,4	9,7	13,6	9,4	10,5	9,8	10,2	15,5	10,6	13,2	10,4	10,9	18,3	11,2	4,00	1,21	1,18	4,86	1,78
T50_HC_PYR	418	420	413	420	425	424	425	419	427	432	431	433	425	432	440	10,3	9,5	10,1	9,4	11,3
T70_HC_PYR	459	461	456	459	465	463	465	460	463	473	468	471	465	467	481	8,4	8,7	7,3	7,0	11,4
T90_HC_PYR	506	512	505	500	523	513	520	510	508	534	523	529	516	515	542	12,7	11,9	9,1	10,5	13,9
T30_CO2_PYR	332	336	331	327	341	337	340	335	332	346	341	345	338	337	353	9,0	7,0	6,8	7,2	11,0
T50_CO2_PYR	377	383	376	371	389	384	389	381	378	396	390	395	387	384	406	11,9	9,7	9,6	9,9	15,4
T70_CO2_PYR	429	435	428	421	441	436	441	434	429	451	443	448	440	437	463	13,3	11,3	10,4	11,9	16,9
T90_CO2_PYR	499	503	500	493	509	505	508	504	500	517	511	514	509	507	525	10,3	8,5	7,7	10,7	11,1
T50_CO_PYR	399	400	398	402	400	403	402	401	406	404	406	405	403	412	408	11,8	11,4	7,8	11,6	7,0
T50_CO_OX	382	388	377	376	403	397	405	391	394	427	413	424	407	405	456	29,4	30,8	24,8	22,6	39,1
T70_CO_OX	426	446	421	418	482	453	475	444	435	506	486	499	475	454	523	39,9	37,8	36,5	32,4	40,6
T50_CO2_OX	403	405	402	403	412	411	413	408	411	427	420	425	416	417	433	13,7	14,5	11,9	11,1	18,0
T70_CO2_OX	444	449	443	439	459	457	468	453	451	491	478	486	466	466	501	22,7	22,8	18,0	21,0	25,9
T90_CO2_OX	518	528	519	503	538	534	540	531	524	551	544	548	539	540	554	20,7	17,5	16,9	23,0	13,9
PseudoS1	0,08	0,06	0,10	0,11	0,06	0,11	0,08	0,13	0,15	0,08	0,16	0,11	0,17	0,22	0,10	0,09	0,04	0,08	0,11	0,04
S2	1,72	1,27	2,73	2,92	0,69	2,99	1,82	3,96	4,47	0,94	5,05	2,88	6,00	7,53	1,46	4,310	1,44	3,96	4,56	0,880
S2_PC	0,6	0,57	0,64	0,64	0,50	0,66	0,62	0,69	0,69	0,53	0,71	0,66	0,73	0,75	0,58	0,08	0,07	0,06	0,08	0,07
HI	174	159	199	192	127	206	183	226	229	137	246	212	257	275	155	57,0	43,5	45,0	66,1	28,9
OIre6	155	169	151	145	175	175	187	168	163	192	195	207	189	181	213	31,5	29,6	26,6	29,7	44,8
HI_OIre6	0,91	0,82	1,09	1,09	0,62	1,17	0,98	1,34	1,39	0,69	1,52	1,21	1,63	1,83	0,86	0,55	0,37	0,43	0,69	0,23
PC	2,90	2,22	4,17	4,26	1,43	4,54	3,04	5,97	6,53	1,99	7,22	4,49	8,63	10,80	2,47	5,59	1,94	5,14	6,09	1,26
PC_TOCre6	0,24	0,23	0,25	0,25	0,21	0,26	0,25	0,27	0,28	0,22	0,29	0,27	0,30	0,31	0,24	0,04	0,03	0,03	0,050	0,029
TOCre6	11,2	9,1	15,3	15,4	6,7	17,0	12,5	21,8	24,3	8,5	26,8	17,1	30,3	38,4	11,4	18,5	7,05	17,5	20,9	4,70
MINC	1,0	0,9	1,1	1,0	0,8	1,6	1,3	1,8	1,7	2,1	4,3	5,2	3,4	3,7	15,9	16,7	19,83	13	14	18,1
R_index	0,57	0,58	0,55	0,58	0,62	0,61	0,61	0,58	0,62	0,65	0,64	0,65	0,61	0,65	0,68	0,05	0,04	0,05	0,05	0,05
I_index	0,08	0,08	0,13	0,05	0,01	0,13	0,13	0,19	0,11	0,09	0,19	0,17	0,23	0,16	0,12	0,08	0,06	0,07	0,08	0,07

695 **Table B: (2 pages) Minimum, maximum, mean, 1st quartile, 3rd quartile, median and standard deviation values of the Rock-Eval® parameters for the RMQS topsoil (0–30 cm) samples limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3 (n=1891, “total”) and by land cover.**



700 **Figure C: Effect of land cover on topsoil organic carbon stoichiometry for the RMQS topsoil (0–30 cm) samples under the four major land covers in France: croplands, forests, grasslands, and vineyards & orchards. (a) C/N distribution; (b) HI distribution;**  
 705 **(c) OIre6 distribution. Samples are limited to those with Rock-Eval® organic carbon yields ranging from 0.7 to 1.3. For boxplots, the black midline of each box is the median. Lower and upper edges are respectively the 1<sup>st</sup> and 3<sup>rd</sup> quartiles. Lower and upper whiskers are respectively the maximum between the minimum value or the 1<sup>st</sup> quartile minus 1.5 times the interquartile range (max[ $\min$ ;  $Q1 - 1.5 \cdot (Q3 - Q1)$ ]) and the minimum between the maximum or the 3<sup>rd</sup> quartile plus 1.5 times the interquartile range (min[ $\max$ ;  $Q3 + 1.5 \cdot (Q3 - Q1)$ ]). Different letters indicate significant differences in the distribution of the values for the land uses with the Kruskal-Wallis test ( $P < 0.05$ ) and multiple Wilcoxon test ( $P < 0.05$ ).**



710 **Figure D:** Score of the 2037 samples on axes 1 and 2 of the principal component analysis on 11 pedoclimatic parameters: clay, total silt and total sand contents, pH in water, water content, carbonate content, mean annual temperature, mean annual precipitation, Tamm and Mehra-Jackson iron oxyhydroxides contents, C/N ratio for the RMQS topsoil (0–30 cm) samples. The samples with an organic carbon yield between 0.7 and 1.3 are plotted in light blue. The samples with an organic carbon yield < 0.7 or > 1.3 are plotted in dark blue.